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Date Complete:	Shorium	Car manch

Author Search

=> FILE HCAPLUS
FILE 'HCAPLUS' ENTERED AT 14:36:23 ON 11 APR 2008
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FILE COVERS 1907 - 11 Apr 2008 VOL 148 ISS 16 FILE LAST UPDATED: 10 Apr 2008 (20080410/ED)

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Structure attributes must be viewed using STN Express query preparation: Uploading strA.str

```
12 13 15 17 19 20
ring nodes :
1 2 3 4 5 6 7 8 9 10
chain bonds :
1-15 2-12 7-13 13-17 19-20
ring bonds :
1-2 1-6 2-3 3-4 4-5 4-7 5-6 5-10 7-8 8-9 9-10
exact/norm bonds :
1-2 \quad 1-6 \quad 1-15 \quad 2-3 \quad 2-12 \quad 3-4 \quad 4-5 \quad 4-7 \quad 5-6 \quad 5-10 \quad 7-8 \quad 7-13 \quad 8-9 \quad 9-10 \quad 13-17
19-20
```

G1:0,S

G2:CH,[*1]

G3:H, X, Ak

Match level :

Saturation

chain nodes :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 12:CLASS 13:CLASS 15:CLASS 17:Atom 19:CLASS 20:CLASS Generic attributes :

17:

: Unsaturated

Element Count : Node 17: Limited N.N1

L3 150 SEA FILE=REGISTRY SSS FUL L1 6 SEA FILE=HCAPLUS ABB=ON PLU=ON L3 L4

L5	6	SEA FILE=HCAPLUS A	ABB=ON PLU=ON	L4 AND (PRY<=2005 OR AY<=2005
		OR PY<=2005)		
L6	138	SEA FILE=HCAPLUS A	ABB=ON PLU=ON	MITSUYA M?/AU
L7	47	SEA FILE=HCAPLUS A	ABB=ON PLU=ON	BAMBA M?/AU
L8	7626	SEA FILE=HCAPLUS A	ABB=ON PLU=ON	SASAKI Y?/AU
L9	6232	SEA FILE=HCAPLUS A	ABB=ON PLU=ON	NISHIMURA T?/AU
L10	19	SEA FILE=HCAPLUS A	ABB=ON PLU=ON	EIKI J?/AU
L11	1742	SEA FILE=HCAPLUS A	ABB=ON PLU=ON	ARAKAWA K?/AU
L12	15764	SEA FILE=HCAPLUS A	ABB=ON PLU=ON	(L6 OR L7 OR L8 OR L9 OR L10
		OR L11)		
L13	1	SEA FILE=HCAPLUS A	ABB=ON PLU=ON	L5 AND L12

=> FILE WPIX

FILE 'WPIX' ENTERED AT 14:36:31 ON 11 APR 2008 COPYRIGHT (C) 2008 THE THOMSON CORPORATION

FILE LAST UPDATED: 8 APR 2008 <20080408/UP>
MOST RECENT THOMSON SCIENTIFIC UPDATE: 200823 / 200823/WD>
DERNENT WORLD PATENTS INDEX SUBSCRIBER FILE, COVERS 1963 TO DATE

>>> IPC Reform backfile reclassification has been loaded to the end of November 2007. No update date (UP) has been created for the reclassified documents, but they can be identified by 20060101/UPIC and 20061231/UPIC, 20070601/UPIC, 20071001/UPIC and 20071130/UPIC. <<</p>

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- >>> XML document distribution format now available See HELP XMLDOC <<<
- >>> ECLA Codes and Current US National Classifications have been added see NEWS and HELP CHANGE <<<</pre>
- >>> HELP for European Patent Classifications see HELP ECLA, HELP ICO <<<
- >>> Updated PDF files in the following links:
 http://www.stn-international.de/stndatabases/details/ico_0803.zip
 http://www.stn-international.de/stndatabases/details/epc_0803.zip
 Supplement of all changed ECLA items:
 http://www.stn-international.de/stndatabases/details/ecla_0803s.zip <<</pre>
- 'BI, ABEX' IS DEFAULT SEARCH FIELD FOR 'WPIX' FILE

=> D QUE L17 L1 STR

```
Structure attributes must be viewed using STN Express query preparation.
           138 SEA FILE=HCAPLUS ABB=ON PLU=ON MITSUYA M?/AU
1.7
            47 SEA FILE=HCAPLUS ABB=ON PLU=ON BAMBA M?/AU
1.8
           7626 SEA FILE-HCAPLUS ABB-ON PLU-ON SASAKI Y?/AU
L9
          6232 SEA FILE=HCAPLUS ABB=ON PLU=ON NISHIMURA T?/AU
1.10
            19 SEA FILE=HCAPLUS ABB=ON PLU=ON EIKI J?/AU
L11
           1742 SEA FILE=HCAPLUS ABB=ON PLU=ON ARAKAWA K?/AU
L12
          15764 SEA FILE=HCAPLUS ABB=ON PLU=ON (L6 OR L7 OR L8 OR L9 OR L10
                OR L11)
1.15
            63 SEA FILE=WPIX SSS FUL L1
L16
             2 SEA FILE=WPIX ABB=ON PLU=ON L15/DCR
L17
              1 SEA FILE=WPIX ABB=ON PLU=ON L12 AND L16
=> DUP REM L13 L17
FILE 'HCAPLUS' ENTERED AT 14:36:40 ON 11 APR 2008
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PROCESSING COMPLETED FOR L13
PROCESSING COMPLETED FOR L17
              1 DUP REM L13 L17 (1 DUPLICATE REMOVED)
                ANSWER '1' FROM FILE HCAPLUS
=> D IBIB ED ABS FHITSTR L22 1
L22 ANSWER 1 OF 1 HCAPLUS COPYRIGHT 2008 ACS on STN DUPLICATE 1
                        2005:1042235 HCAPLUS Full-text
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         143:347192
TITLE:
                         Preparation of substituted quinazoline and
                         pyridopyrimidine derivatives as glucokinase activators
INVENTOR(S):
                         Missuva, Morihiro; Bamba, Makoto;
                         Sasaki, Yasuhiro: Nishimura, Teruvuki
                         ; Eiki, Junichi; Arakawa, Keisuhe
PATENT ASSIGNEE(S):
                         Banyu Pharmaceutical Co., Ltd, Japan
                        PCT Int. Appl., 192 pp.
SOURCE:
```

CODEN: PIXXD2 Patent

Japanese

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.						APPLICATION NO.												
WO	2005	0903	32		A1				WO 2005-JP5991								<	
	W:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FΙ,	GB,	GD,	
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NA,	NI,	
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	
		SY,	TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
	RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑM,	
		AZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	
		EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,	IS,	IT,	LT,	LU,	MC,	NL,	PL,	PT,	
		RO,	SE,	SI,	SK,	TR,	BF,	ΒJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	
		MR,	NE,	SN,	TD,	TG												
AU	2005	2236	10		A1		2005	0929		AU 2	005-	2236	10		2	0050	323	<
CA	2560	286			A1		2005	0929		CA 2	005-	2560	286		2	0050	323	<
EP	1734	040			A1		2006	1220		EP 2	005-	7216	40		2	0050	323	<
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		IS,	IT,	LI,	LT,	LU,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	LV		
CN	1934	100			A		2007	0321		CN 2	005-	8000	9447		2	0050	323	<
IN	2006	DN05	151		A		2007	0803		IN 2	006-	DN51	51		2	0060	907	<
US	2008	0032	996		A1		2008	0207		US 2	007-	5935	40		2	0070	510	<
IORIT	Y APP	LN.	INFO	. :						JP 2	004-	8580	8		A 2	0040	323	<
										WO 2	005-	JP59	91		W 2	0050	323	<
HER S	OURCE	(S):			MAR	PAT	143:	3471	92									
En:	tered	STN	: 2	9 Se	p 20	05												

AB The title compds. I [X is a nitrogen atom, CH; Y is O, S; R1 is an optionally substituted 5 to 6-membered heteroaryl group, aryl, alkyl, etc.; R2 is a hydrogen atom or a fluorine atom; and the ring A is an optionally substituted monocyclic or bicyclic heteroaryl group] are prepared Thus, [6-(4H-[1,2,4]triazol-3-ylsulfanyl)quinazolin-4-yl]thiazolo[5,4-b]pyridin-2-ylamine was prepared in 2 steps from 4-chloro-6-iodoquinazoline and thiazolo[5,4b]pyridin-2-ylamine. In a test for glucokinase activating activity, compds. of this invention showed EC50 values of 0.08 µM to 0.18 µM. Formulations are given.

ΙT 865662-62-0P RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

> (preparation of substituted quinazoline and pyridopyrimidine derivs. as glucokinase activators)

RN 865662-62-0 HCAPLUS

CN 6-Quinazolinol, 4-(thiazolo[5,4-b]pyridin-2-ylamino)- (CA INDEX NAME)



ONE OR MORE TAUTOMERIC DOUBLE BONDS NOT DISPLAYED IN THE STRUCTURE
REFERENCE COUNT: 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS

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Structure Search

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FILE COVERS 1907 - 11 Apr 2008 VOL 148 ISS 16 FILE LAST UPDATED: 10 Apr 2008 (20080410/ED)

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=> D QUE L5

L1 STR

Structure attributes must be viewed using STN Express query preparation.

L3 150 SEA FILE=REGISTRY SSS FUL L1

L4 6 SEA FILE=HCAPLUS ABB=ON PLU=ON L3

L5 6 SEA FILE-HCAPLUS ABB-ON PLU-ON L4 AND (PRY<-2005 OR AY<-2005

OR PY<=2005)

=> S L5 NOT L13

L23 5 L5 NOT L13

=> FILE WPIX

FILE 'WPIX' ENTERED AT 14:37:14 ON 11 APR 2008 COPYRIGHT (C) 2008 THE THOMSON CORPORATION

FILE LAST UPDATED: 8 APR 2008 <20080408/UP>
MOST RECENT THOMSON SCIENTIFIC UPDATE: 200823 <200823/DW>
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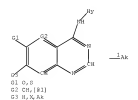
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- >>> XML document distribution format now available See HELP XMLDOC <<<
- >>> ECLA Codes and Current US National Classifications have been added see NEWS and HELP CHANGE <<<
- >>> HELP for European Patent Classifications see HELP ECLA, HELP ICO <<<
- >>> Updated PDF files in the following links: http://www.stn-international.de/stndatabases/details/ico_0803.zip http://www.stn-international.de/stndatabases/details/epc_0803.zip Supplement of all changed ECLA items:
- http://www.stn-international.de/stndatabases/details/ecla_0803s.zip <<< 'BI,ABEX' IS DEFAULT SEARCH FIELD FOR 'WPIX' FILE



Structure attributes must be viewed using STN Express query preparation.

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Serial No.:10/593,540
L15
          63 SEA FILE-WPIX SSS FUL L1
L16
            2 SEA FILE-WPIX ABB-ON PLU-ON L15/DCR
=> S L16 NOT L17
L24 1 L16 NOT L17
=> FILE BABS
FILE 'BABS' ENTERED AT 14:37:34 ON 11 APR 2008
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FILE LAST UPDATED: 17 MAR 2008 <20080317/UP>
FILE COVERS 1980 TO DATE
=> D OUE L20
1.20
            1 SEA FILE=BABS ABB=ON PLU=ON 6424720/BABSAN
=> FILE BEILSTEIN
FILE 'BEILSTEIN' ENTERED AT 14:37:44 ON 11 APR 2008
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FILE LAST UPDATED ON April 1, 2008
FILE COVERS 1771 TO 2008.
*** FILE CONTAINS 10.322,808 SUBSTANCES ***
>>>PLEASE NOTE: Reaction Data and substance data are stored in
  separate documents and can not be searched together in one query.
  Reaction data for BEILSTEIN compounds may be displayed
  immediately with the display codes PRE (preparations) and REA
  (reactions). A substance answer set retrieved after the search
  for a chemical name, a compounds with available reaction
  information by combining with PRE/FA, REA/FA or more generally
  with RX/FA. The BEILSTEIN Registry Number (BRN) is the link
  between a BEILSTEIN compound and belonging reactions. For mo
  detailed reaction searches BRNs can be searched as reaction
  partner BRNs Reactant BRN (RX.RBRN) or Product BRN (RX.PBRN). <<<
>>> FOR SEARCHING PREPARATIONS SEE HELP PRE <<<
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>>> Price change as of January 1st, 2008: Connect Time and Structure Search fees re-introduced. See NEWS and HELP COST <<<

=> D QUE L21 STR T.1

Structure attributes must be viewed using STN Express query preparation. T.18 4 SEA FILE=BEILSTEIN SSS FUL L1

1.19 1 SEA FILE=BEILSTEIN ABB=ON PLU=ON L18 AND BABSAN/FA

1.21 3 SEA FILE=BEILSTEIN ABB=ON PLU=ON L18 NOT L19

=> DUP REM L23 L24 L20 L21

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ANSWERS FROM THESE FILES WILL BE CONSIDERED UNIQUE FILE 'HCAPLUS' ENTERED AT 14:37:59 ON 11 APR 2008

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PROCESSING COMPLETED FOR L23

PROCESSING COMPLETED FOR L24

PROCESSING COMPLETED FOR L20

PROCESSING COMPLETED FOR L21

L25 8 DUP REM L23 L24 L20 L21 (2 DUPLICATES REMOVED) ANSWERS '1-5' FROM FILE HCAPLUS

ANSWERS '6-8' FROM FILE BEILSTEIN

=> D IBIB ED ABS HITSTR 1-5; D IDE ALLREF 6-8

L25 ANSWER 1 OF 8 HCAPLUS COPYRIGHT 2008 ACS on STN DUPLICATE 1

ACCESSION NUMBER: 2003:981459 HCAPLUS Full-text

DOCUMENT NUMBER: 140:228690

TITLE: Synthesis and SAR of potent EGFR/erbB2 dual inhibitors AUTHOR(S): Zhang, Yue-Mei; Cockerill, Stuart; Guntrip, Stephen B.; Rusnak, David; Smith, Kathryn; Vanderwall, Dana;

Wood, Edgar; Lackey, Karen

CORPORATE SOURCE: GlaxoSmithKline, Research Triangle Park, NC, 27709,

IISA

SOURCE: Bioorganic & Medicinal Chemistry Letters (2004

), 14(1), 111-114

CODEN: BMCLE8; ISSN: 0960-894X

Elsevier Science B.V. PUBLISHER:

DOCUMENT TYPE: Journal LANGUAGE: English

Entered STN: 17 Dec 2003 ED

AΒ A series of 6-alkoxy-4-anilinoquinazoline compds, was prepared and evaluated for in vitro inhibition of the erbB2 and EGFR kinase activity. The IC50 values of the best compds. were below 0.10 uM. Further, several of these

compds, inhibit the growth of erbB2 and EGFR over-expressing tumor cell lines at concns. below 1 uM.

668437-13-6P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(synthesis and structure-activity relations of potent EGFR/erbB2 kinase dual inhibitors)

668437-13-6 HCAPLUS

CN 4-Quinazolinamine, N-[1-[(3-fluorophenyl)methyl]-1H-indazol-5-yl]-6-[4-[[(methylsulfonyl)methyl]amino]butoxy]- (CA INDEX NAME)

REFERENCE COUNT: 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L25 ANSWER 2 OF 8 HCAPLUS COPYRIGHT 2008 ACS on STN DUPLICATE 2

ACCESSION NUMBER: 1999:451283 HCAPLUS Full-text

DOCUMENT NUMBER: 131:102287

TITLE: Preparation of quinazolinvlamines and analogs as

protein tyrosine kinase inhibitors

INVENTOR(S): Cockerill, George Stuart; Lackey, Karen Elizabeth

PATENT ASSIGNEE(S): Glaxo Group Limited, UK

SOURCE: PCT Int. Appl., 145 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent. LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE 19990715 WO 9935132 A1 WO 1999-GB76 19990111 <--W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP,

KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM,

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TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

AU 9919786 A 19990726 AU 1999-19786 19990111 <---
PRIORITY APPLN. INFO:: GB 1998-575 A 19980112 <---
WO 1999-GB76 W 19990111 <---
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OTHER SOURCE(S): MARPAT 131:102287
ED Entered STN: 23 Jul 1999
GI

AB Substituted heteroarom. compds. I are prepared [wherein X = N or CH; Y = CR1 and V = N; or Y = N and V = CR1; or Y = CR1 and V = CR2; or Y = CR2 and V = CR1: R1 = O-M-, wherein M = C1-5 alkylene where any C atom not immediately adjacent to Q may be replaced by O, S, or NR6; Q = wide variety of groups; R2 = H, halo, OH, alkyl, alkoxy, (di)alkylamino; U = Ph, pyridyl, pyrimidinyl, imidazolyl, or 9- or 10-membered bicyclic heterocyclyl containing 1-2 N atoms and 0-1 addnl. O, N, or S; U is substituted by R3, where R3 = benzyl, halobenzyl, pyridylmethyl, pyridylmethoxy, PhO, PhSO2, (un)substituted phthalimido; R6 = H, alkvll. Twelve examples and a variety of intermediates were prepared For instance, 4-chloro-6-iodoquinazoline was aminated in the 4position with 5-amino-1-benzyl-1H-indazole, followed by Pd-catalyzed carbonylation, to give 4-[(1-benzyl-1H-indazol-5-yl)amino]quinazoline-6carbaldehyde. This underwent reductive amination by MeSO2CH2CH2NH2 and a reducing agent such as NaBH(OAc)3, to give title compound II.HCl. In an EGFr phosphorylation assay, II.HCl had an IC50 of <0.10 uM. IΤ 230955-59-6P 230955-69-9P 230955-73-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; preparation of quinazolinylamines and analogs as protein tyrosine kinase inhibitors)

RN 230955-59-6 HCAPLUS

CN 6-Quinazolinol, 4-[[1-[(3-fluorophenyl)methyl]-1H-indazol-5-yl]amino]-, acetate (ester) (9CI) (CA INDEX NAME)

- RN 230955-60-9 HCAPLUS
- CN 6-Quinazolinol, 4-[[1-[(3-fluorophenyl)methyl]-1H-indazol-5-yl]amino]-(CA INDEX NAME)

- RN 230955-73-4 HCAPLUS
- CN Acetamide, 2,2,2-trifluoro-N-[4-[[4-[[1-[3-fluorophenyl]methyl]-1H-indazol-5-yl]amino]-6-quinazolinyl]oxy]butyl]-N-[2-(methylsulfonyl)ethyl]-(CA INDEX NAME)

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- IT 230955-49-4P
 - RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 - (target compound; preparation of quinazolinylamines and analogs as protein tyrosine kinase inhibitors)
- RN 230955-49-4 HCAPLUS
- CN 4-Quinazolinamine, N-[1-[(3-fluoropheny1)methy1]-1H-indazol-5-y1]-6-[4-[[2-(methylsulfony1)ethy1]amino]butoxy]- (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

REFERENCE COUNT:

→ F

SOURCE:

6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L25 ANSWER 3 OF 8 HCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2001:743253 HCAPLUS Full-text

DOCUMENT NUMBER: 136:79264

TITLE: The characterization of novel, dual ErbB-2/EGFR,

tyrosine kinase inhibitors: potential therapy for

cancer

AUTHOR(S): Rusnak, David W.; Affleck, Karen; Cockerill, Stuart G.; Stubberfield, Colin; Harris, Robert; Page, Martin;

Smith, Kathryn J.; Guntrip, Stephen B.; Carter, Malcolm C.; Shaw, Robert J.; Jowett, Amanda; Stables, Jeremy; Topley, Peter; Wood, Edgar R.; Brignola, Perry S.; Kadwell, Sue H.; Reep, Bryan R.; Mullin, Robert J.; Alligood, Krystal J.; Keith, Barry R.; Crosby,

Renae M.; Murray, Doris M.; Knight, W. Blaine; Gilmer,

Tona M.; Lackey, Karen CORPORATE SOURCE:

Department of Cancer Biology, GlaxoSmithKline, Research Triangle Park, NC, 27709, USA

Cancer Research (2001), 61(19), 7196-7203 CODEN: CNREA8; ISSN: 0008-5472

PUBLISHER: American Association for Cancer Research DOCUMENT TYPE: Journal

LANGUAGE: English ED Entered STN: 11 Oct 2001

AB The type 1 receptor tyrosine kinases constitute a family of transmembrane proteins involved in various aspects of cell growth and survival and have been implicated in the initiation and progression of several types of human malignancies. The best characterized of these proteins are the epidermal growth factor receptor (EGFR) and ErbB-2 (HER-2/neu). We have developed potent quinazoline and pyrido-[3,4-d]-pyrimidine small mols, that are dual inhibitors of ErbB-2 and EGFR. The compds. demonstrate potent in vitro

inhibition of the ErbB-2 and EGFR kinase domains with IC50s <80 nM. Growth of

ErbB-2- and EGFR-expressing tumor cell lines is inhibited at concns. <0.5 μM. Selectivity for tumor cell growth inhibition vs. normal human fibroblast growth inhibition ranges from 10- to >75-fold. Tumor growth in mouse s.c. xenograft models of the BT474 and Rh5 cell lines is inhibited in a doseresponsive manner using oral doses of 10 and 30 mg/kg twice per day. In addition, the tested compds. caused a reduction of ErbB-2 and EGFR autophosphorylation in tumor fragments from these xenograft models. These data indicate that these compds. have potential use as therapy in the broad population of cancer patients overexpressing ErbB-2 and/or EGFR. 220955-49-4, GW 5945

RL: PAC (Pharmacological activity); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(characterization of novel, dual ErbB-2/EGFR, tyrosine kinase inhibitors and potential therapy for cancer)

RN 230955-49-4 HCAPLUS

TT

CN 4-Quinazolinamine, N-[1-[(3-fluorophenyl)methyl]-1H-indazol-5-yl]-6-[4-[[2-(methylsulfonyl)ethyl]amino]butoxy]- (CA INDEX NAME)

PAGE 1-B

REFERENCE COUNT: 48 THERE ARE 48 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L25 ANSWER 4 OF 8 HCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1999:113672 HCAPLUS Full-text

DOCUMENT NUMBER: 130:182476

TITLE: Preparation of heterocyclic compounds as irreversible

bicyclic inhibitors of tyrosine kinases

INVENTOR(S): Bridges, Alexander James

PATENT ASSIGNEE(S): Warner-Lambert Company, USA SOURCE: PCT Int. Appl., 131 pp.

DOCUMENT TYPE: CODEN: PIXXD2
Patent

DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.			KIND DATE			APPLICATION NO.											
WO	9906	396							WO 1998-US15592					19980729 <				
	W:	AL,	AU,	BA,	BB,	BG,	BR,	CA,	CN,	CZ	, EE,	GE,	HR,	HU,	ID,	IL,	IS,	
		JP,	KR,	LC,	LK,	LR,	LT,	LV,	MG,	MK	, MN,	MX,	NO,	NZ,	PL,	RO,	SG,	
		SI,	SK,	SL,	TR,	TT,	UA,	US,	UZ,	VN	, YU,	AM,	ΑZ,	BY,	KG,	ΚZ,	MD,	
		RU,	ТJ,	TM														
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		FI,	FR,	GB,	GR,	IE,	ΙT,	LU,	MC,	NL	, PT,	SE,	BF,	ΒJ,	CF,	CG,	CI,	
		CM,	GΑ,	GN,	GW,	ML,	MR,	NE,	SN,	TD	, TG							
AU	9886	659			A		1999	0222		AU	1998-	8665	9		1	9980	729	<
US	6153	617			A		20001128			US 1999-269647			19990325 <					
US	2003	0087	881		A1		2003	0508		US	2002-	2726.	51		2	0021	017	<
PRIORIT:	Y APP	LN.	INFO	.:						US	1997-	5406	1P		P 1	9970	729	<
										WO	1998-	US15.	592		W 1	9980	729	<
										US	1999-	2696	47		A3 1	9990	325	<
										US	2000-	6563	31		B1 2	0000	906	<
OTHER SO	DURCE	(S):			MAR	PAT	130:	1824	76									

X N Z

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ED Entered STN: 19 Feb 1999

AB The title compds., e.g. I [X = DEF, Y = SR4, etc.; or X = SR4, etc., and Y = DEF; D = O, etc.; E = CO, etc.; F = CR1(:C):C(R5)H, etc.; a proviso is given; R1 = H, halo, etc.; R5 = H, halo, perfluoroalkyl, etc.; Z = indoline molety (generic structure given), etc.; R4 = H, alkyl, etc.], are prepared This invention also provides a method of treating cancer, restenosis, atherosclerosis, endometriosis, and psoriasis and a pharmaceutical composition that comprises a compound that is an irreversible inhibitor of tyrosine kinases. N-[4-(6-bromo-2,3-dihydroindol-1- yl)quinazolin-6-yl)acrylamide in vitro showed IC50 of 0.4 nM against epidermal growth factor receptor tyrosine kinase.

IT 220577-65-1P 220577-66-2P 220577-73-1P 220577-74-2P 220577-75-3P 220577-76-4P 220577-75-9 220577-79-7P 220577-82-2P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of heterocyclic compds. as irreversible bicyclic inhibitors of tyrosine ${\tt kinases})$

220577-65-1 HCAPLUS

CN 2-Propenoic acid, 4-(1H-benzimidazol-5-ylamino)-6-quinazolinyl ester (9CI) (CA INDEX NAME)

RN 220577-66-2 HCAPLUS

CN 2-Propenoic acid, 4-(1H-indol-5-ylamino)-6-quinazolinyl ester (CA INDEX NAME)

RN 220577-73-1 HCAPLUS

CN 2-Octenoic acid, 4,4-difluoro-8-(4-morpholinyl)-, 4-[(1-methyl-1H-indol-5-yl)amino]-6-quinazolinyl ester (CA INDEX NAME)

RN 220577-74-2 HCAPLUS

CN 2-Octenoic acid, 4,4-difluoro-8-(4-morpholiny1)-, 4-(1H-indol-5-ylamino)-6quinazoliny1 ester (CA INDEX NAME)

CN 2-Pentenoic acid, 5-[[3-(4-morpholinyl)propyl]amino]-5-oxo-, 4-(1H-indol-5-ylamino)-6-quinazolinyl ester (CA INDEX NAME)

RN 220577-76-4 HCAPLUS

CN 2-Pentenoic acid, 5-[[3-(4-morpholiny1)propy1]amino]-5-oxo-,
4-[(1-methyl-1H-indol-5-yl)amino]-6-quinazoliny1 ester (CA INDEX NAME)

RN 220577-77-5 HCAPLUS

CN 2-Butenoic acid, 4-[[3-(4-morpholiny1)propy1]thio]-, 4-[(2-methyl-1H-indol-5-yl)amino]-6-quinazolinyl ester (CA INDEX NAME)

RN 220577-78-6 HCAPLUS

CN 2-Butenoic acid, 4-[[3-(4-morpholinyl)propyl]thio]-, 4-(1H-indol-5ylamino)-6-quinazolinyl ester (CA INDEX NAME)

RN 220577-79-7 HCAPLUS

CN 2-Heptenoic acid, 7-(4-morpholinyl)-, 4-[(3-cyano-1H-indol-5-yl)amino]-6quinazolinyl ester (CA INDEX NAME)

RN 220577-80-0 HCAPLUS

CN 2-Heptenoic acid, 7-(4-morpholinyl)-, 4-(1H-indol-5-ylamino)-6quinazolinyl ester (CA INDEX NAME)

RN 220577-82-2 HCAPLUS

CN 2-Butenoic acid, 4-(4-morpholinyl)-, 4-(1H-indol-5-ylamino)-6-quinazolinyl ester (CA INDEX NAME)

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L25 ANSWER 5 OF 8 HCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1995:23238 HCAPLUS Full-text

DOCUMENT NUMBER: 122:31545

TITLE: Preparation of aminoquinazolines useful in the

treatment of cancer

INVENTOR(S): Barker, Andrew John; Brown, Dearg Sutherland

PATENT ASSIGNEE(S): Zeneca, UK

SOURCE: Eur. Pat. Appl., 39 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.		D DATE	APPLICATION NO.	
	P 602851 P 602851	A1 B1		EP 1993-309680	19931203 <
	R: AT, BE	, CH, DE,	DK, ES, FR,	GB, GR, IE, IT, LI,	LU, MC, NL, PT, SE
F	U 9350728	A	19940623	AU 1993-50728	19931116 <
F	U 664496	B2	19951116		
2	A 9308594	A	19940610	ZA 1993-8594	19931117 <
0	A 2103383	A1	19940611	CA 1993-2103383	19931118 <
0	A 2103383	C	20050125		
]	L 107678	A	19990312	IL 1993-107678	19931119 <
F	U 65622	A2	19940728	HU 1993-3328	19931124 <
E	'I 9305431	A	19940611	FI 1993-5431	19931203 <
P	T 143956	T	19961015	AT 1993-309680	19931203 <
E	S 2093367	Т3	19961216	ES 1993-309680	19931203 <
0	Z 283612	B6	19980513	CZ 1993-2651	19931206 <
N	0 9304504	A	19940613	NO 1993-4504	19931209 <
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Ċ	P 3330706	B2	20020930		
0	N 1094043	A	19941026	CN 1993-120872	19931210 <
Ţ	IS 5580870	A	19961203	US 1993-164725	19931210 <
PRIORI	TY APPLN. INFO	O.:		GB 1992-25765	A 19921210 <
				GB 1993-10248	A 19930518 <

OTHER SOURCE(S): MARPAT 122:31545

ED Entered STN: 08 Nov 1994

GI

- AB The title compds. [I; Q = 9- or 10-membered bicyclic heterocyclic moiety containing 1-2 N atoms; R1 = OH, NH2, ureido, hydroxyamino, trifluoromethoxy, (un)substituted C1-4 alkyl, C1-4 alkoxy, pyrrolidin-1-yl, piperidino, etc.; m = 1-3], useful in the treatment of cancer (no data), are prepared and I-containing formulations presented. Thus, 4-chloro-6,7-dimethoxyquinazoline was reacted with 5-aminoquinoline, producing 6,7-dimethoxy-4-(5-quinolylamino)quinazoline, m.p. > 240° in 35% yield.
- IT 159768-49-7P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SFN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of, as anticancer agent)

RN 159768-49-7 HCAPLUS

CN 4-Quinazolinamine, 7-fluoro-N-1H-indol-5-yl-6-methoxy-, monohydrochloride (9CI) (CA INDEX NAME)

Molec. Formula (MF):

L25 ANSWER 6 OF 8 BEILSTEIN COPYRIGHT 2008 BEILSTEIN MDL on STN

Beilstein Records (BRN): 9602955
Chemical Name (CN): 2,2,2-trifluoro-N-(4-<4-<1-(3-fluoro-benzyl)-1H-indazol-5-ylamino>-quinazolin-6-

yloxy>-butyl)-N-(2-methanesulfonyl-ethyl)acetamide

Autonom Name (AUN): 2.2.2-trifluoro-N-(4-<4-<1-(3-fluoro-

benzyl)-1H-indazol-5-ylamino>-quinazolin-6yloxy>-butyl)-N-(2-methanesulfonyl-ethyl)-

acetamide C31 H30 F4 N6 O4 S

Molecular Weight (MW): 658.67

Lawson Number (LN): 29684, 29566, 16445, 3140, 3125, 1157, 292

Compound Type (CTYPE): heterocyclic Constitution ID (CONSID): 8097768
Tautomer ID (TAUTID): 9008192
Entry Date (DED): 2004/04/23
Update Date (DUPD): 2004/04/23

Field Availability:

Code Name Occurrence

BRN	Beilstein Records	1
CN	Chemical Name	1
AUN	Autonomname	1
MF	Molecular Formula	1
FW	Formular Weight	1
LN	Lawson Number	7
CTYPE	Compound Type	1
CONSID	Constitution ID	1
TAUTID	Tautomer ID	1
ED	Entry Date	1
UPD	Update Date	1

This substance also occurs in Reaction Documents:

Code	Name	Occurrence
RX	Reaction Documents	2
RXREA	Substance is Reaction Reactant	1
RXPRO	Substance is Reaction Product	1

All References:

ALLREF

 Zhang, Yue-Mei; Cockerill, Stuart; Guntrip, Stephen B.; Rusnak, David; Smith, Kathryn; Vanderwall, Dana; Wood, Edgar; Lackey, Karen, Bioorg.Med.Chem.Lett., CODEN: BMCLE8, 14(1), <2004>, 111 - 114; BABS-6424720

L25 ANSWER 7 OF 8 BEILSTEIN COPYRIGHT 2008 BEILSTEIN MDL on STN

9589052 Beilstein Records (BRN): acetic acid 4-<1-(3-fluoro-benzyl)-1H-Chemical Name (CN): indazol-5-ylamino>-quinazolin-6-yl ester acetic acid 4-<1-(3-fluoro-benzyl)-1H-Autonom Name (AUN): indazol-5-vlamino>-quinazolin-6-vl ester Molec. Formula (MF): C24 H18 F N5 O2 Molecular Weight (MW): 427.44 Lawson Number (LN): Compound Type (CTYPE): Constitution ID (CONSID): 29684, 29566, 16445, 1155 heterocyclic 8086336 Tautomer ID (TAUTID): 9002439 Entry Date (DED): Update Date (DUPD): 2004/04/23 2004/04/23

Field Availability:

Code	Name	Occurrence
BRN	Beilstein Records	1
CN	Chemical Name	1
AUN	Autonomname	1
MF	Molecular Formula	1
FW	Formular Weight	1
LN	Lawson Number	4
CTYPE	Compound Type	1
CONSID	Constitution ID	1
TAUTID	Tautomer ID	1
ED	Entry Date	1
UPD	Update Date	1

This substance also occurs in Reaction Documents:

Code	Name	Occurrence
RX	Reaction Documents	2
RXREA	Substance is Reaction Reactant	1
RXPRO	Substance is Reaction Product	1

All References:

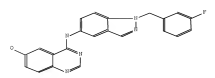
ALLREF

 Zhang, Yue-Mei; Cockerill, Stuart; Guntrip, Stephen B.; Rusnak, David; Smith, Kathryn; Vanderwall, Dana; Wood, Edgar; Lackey, Karen, Bioorg.Med.Chem.Lett., CODEN: BMCLE8, 14(1), <2004>, 111 - 114; BABS-6424720

L25 ANSWER 8 OF 8 BEILSTEIN COPYRIGHT 2008 BEILSTEIN MDL on STN

Beilstein Records (BRN): 9584302 Chemical Name (CN): 4-<1-(3-fluoro-benzyl)-1H-indazol-5ylamino>-quinazolin-6-ol Autonom Name (AUN): 4-<1-(3-fluoro-benzyl)-1H-indazol-5ylamino>-quinazolin-6-ol Molec. Formula (MF): C22 H16 F N5 O Molecular Weight (MW): 385.40 Lawson Number (LN): 29684, 29566, 16445 Compound Type (CTYPE): heterocyclic Constitution ID (CONSID): 8082402

Tautomer ID (TAUTID): 8997062 Entry Date (DED): 2004/04/23 Update Date (DUPD): 2004/04/23



Field Availability:

Code	Name	Occurrence
BRN	Beilstein Records	1
CN	Chemical Name	1
AUN	Autonomname	1
MF	Molecular Formula	1
FW	Formular Weight	1
LN	Lawson Number	3
FS	File Segment	1
CTYPE	Compound Type	1
CONSID	Constitution ID	1
TAUTID	Tautomer ID	1
ED	Entry Date	1
UPD	Update Date	1

This substance also occurs in Reaction Documents:

Code	Name Oc	currence
RX	Reaction Documents	2
RXREA	Substance is Reaction Reactant	1
RXPRO	Substance is Reaction Product	1

All References:

ALLREF

 Zhang, Yue-Mei; Cockerill, Stuart; Guntrip, Stephen B.; Rusnak, David; Smith, Kathryn; Vanderwall, Dana; Wood, Edgar; Lackey, Karen, Bioorg.Med.Chem.Lett., CODEN: BMCLE8, 14(1), <2004>, 111 - 114; BABS-6424720

Search History

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          150 SEA SSS FUL L1
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            6 SEA ABB=ON PLU=ON L4 AND (PRY<=2005 OR AY<=2005 OR PY<=2005)
L5
          138 SEA ABB=ON PLU=ON MITSUYA M?/AU
L6
L7
           47 SEA ABB=ON PLU=ON BAMBA M?/AU
         7626 SEA ABB=ON PLU=ON SASAKI Y?/AU
L8
L9
         6232 SEA ABB=ON PLU=ON NISHIMURA T?/AU
L10
           19 SEA ABB=ON PLU=ON EIKI J?/AU
         1742 SEA ABB=ON PLU=ON ARAKAWA K?/AU
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L12
         15764 SEA ABB=ON PLU=ON (L6 OR L7 OR L8 OR L9 OR L10 OR L11)
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             1 SEA ABB=ON PLU=ON L5 AND L12
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L15
           63 SEA SSS FUL L1
            2 SEA ABB=ON PLU=ON L15/DCR
1.16
            1 SEA ABB=ON PLU=ON L12 AND L16
L17
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              SEL BABSAN
             1 SEA ABB=ON PLU=ON L18 AND BABSAN/FA
L19
              SEL BABSAN
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            1 SEA ABB=ON PLU=ON 6424720/BABSAN
    FILE 'BEILSTEIN' ENTERED AT 14:35:08 ON 11 APR 2008
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L22
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    FILE 'HCAPLUS' ENTERED AT 14:36:58 ON 11 APR 2008
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1.23
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L24
          1 SEA ABB=ON PLU=ON L16 NOT L17
   FILE 'HCAPLUS, WPIX, BABS, BEILSTEIN' ENTERED AT 14:37:59 ON 11 APR 2008
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8 DUP REM L23 L24 L20 L21 (2 DUPLICATES REMOVED)